DEVELOPMENT OF PARTIALLY STABILIZED ZIRCONIA FOR MAKING CERAMIC BUSHINGS

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By
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DEDICATED

TO THE DIVINE SOUL OF MY DEAREST FATHER WHO WAS THE SOURCE OF ALL INSPIRATIONS TO ME.

I PAY TRIBUTES TO MY

DECEASED MOTHER

CERTIFICATE

This is to certify that this work on "Development of Partially stabilized Zirconia for making ceramic bushings" by Narayan Chandra Biswas has been carried out under my supervision and that this has not been submitted elsewhere for a degree.

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SYNOPSIS

The present investigation involved the study of titan addition on Calcia-Zirconia systems in respect to mechanical, thermal and corrosion characteristics. The optimum compositic in respect to bushing material for glass fiber drawing has been searched for. Compositions like 7% (mole) CaO-ZrO₂, 10% CaO-ZrO₂, 13% CaO-ZrO₂, 15% CaO-ZrO₂, 8% MgO-ZrO₂, 7% CaO-5% TiO₂-ZrO₂, 8% CaO-7% TiO₂-ZrO₂, 7% TiO₂-ZrO₂ have been tried using final sintering temperature of 1950°C. Modulus of Rupture, toughness and hardness tests have been done for mechanical characterization.

Vickers hardness value of CaO-ZrO $_2$ and CaO-TiO $_2$ compositions are comparable. 15% CaO-ZrO $_2$ composition has the highest VPH number.

Spalling characteristics is better for CaO-TiO $_2$ -ZrO batches. The composition 7% CaO-5% TiO $_2$ -ZrO $_2$ is best from the spalling resistance point of view.

Corrosion test has been done on sintered crucibles in contact with E glass for 13 hours at 1400°C. Corrosion resistance is satisfactory for titania—added compositions.

CHAPTER-I

INTRODUCTION

Zirconia is a focus of considerable current interest as an outstanding high temperature oxide outweighing the conventional refractories, as a solid electrolyte, and is a unique field of research in consideration to its stabilization, defect crystallography, toughening and strengthening & electrical transport properties etc.

(1,2). very high corrosion resistance to metals, slags and glasses it is used in such applications where temperature and corrosion are very high. Such applications include high temperature extrusion dies, continuous steel pouring nozzles(3), high temperature furnace lining upto 2400°C, lining of furnaces for metting Al, Cr, Co, Au, Ir, Ni, Pd, Pt, Rh metals(4) thermal barrier coatings in air craft and missile components (5), solid electrolyte in fuel cells and steel melts (6), abrasives for melting applications. Glass fibres an usually drawn through bushings made of platinum-rhodium. The high cost of platinum and rhodium makes such process capital-intensive. Recently, ceramic bushings made of alumina as well as stabilised zirconia have been used for making glass fibres (7).

Al202 is a cheaper material, but does not withstand the corrosive effect of E-glass. Zirconia seems to be a solution to this problem. But zirconia has three polymerphs (2). The stable room temperature form is monoclinic. On heating to about 1100°C it transforms to a tetragonal structure which on further heating above 2300°C transforms to a fluorite type cubic structure. Following the works of Murray, Alison, Wolten, Baily and Patil; Bansal and Heuer (8) have conclusively proved that the monoclinic-tetragonal transformation has all the characteristics of martensitic transformations, i.e. it is of an athermal type, apparently diffusionless & involves a large hysteresis (fig. 1.1) (9). The transformation (forward) on cooling occurs over a much lower temperature range (950-850°C) than the reverse transformation (1100-1190°C) on heating. monoclinic-tetragonal phase transition renders the material useless as a high temperature structural material, as it is associated with a large volume change (about 9% volume increase on cooling) which cause cracking and subsequent failure of fabricated products. The transformation interval is displaced in the direction of the lower temperature (10) by addition of metallic oxides with a heterapolar bond energy. phase changes may be suppressed by stabilization of the high temperature cubic fluorite form (fig. 1.1) by alloying with

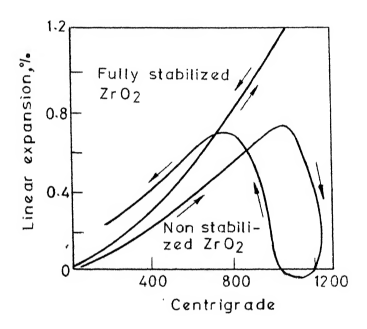


FIG.1.1 THERMAL EXPANSION OF ZIRCONIA MODIFICATIONS (9)

oxides such as those of Mg, Ca, or yttrium. The cubic zirconia solid solution (cubic SS) is infact metastable in all the three systems; alloys of cubic $2rO_2$ with MgO, CaO or Y_2O_3 . The cubic SS tend to decompose by eutectoid decomposition as predicted from phase equilibria considerations, (fig. 2b). In addition, to the tendency to destabilize, fully stabilized $2rO_2$ ceramics have poor thermal shock resistance, because of a combination of low thermal conductivity and high thermal expansion.

It is generally accepted, however, that the most useful mechanical properties are obtained by partial stabilization (11-16) so that a two or three phase microstructure results. These materials are referred to as partially stabilized zirconia (psz). Compositions lying within the cubic field at the firing temperatures and producing monophase microstructure are "fully stabilized" (15).

In recent years, it has been shown that useful mechanical properties of psz, i.e., good fracture toughness and thermal shock resistance can be attributed to a fine dispersion of monoclinic precipitates in the cubic grains (14-15, 17-20). These coherent precipitates are thought to impede crack propagation due to the resulting compressive stresses present in the matrix caused by martensitic

transformation in the particles. The strengthening observed in psz results from crack branching caused by interaction between the stress field of the propagatig crack and residual stress fields around the coherent particles.

If the precipitate size be larger than 2-6 micron, then extensive microcracking takes place due to volume changes on cooling. Because of their large numbers, these cracks propagate only quasistatically and the body maintains a larger portions of its strength after continuous thermal cycling, thus resulting in high thermal shock resistance.

More recently the existence of tetragonal zirconia in sintered bodies with high strength and fracture toughness have been reported (11-13, 16, 21-24). The stabilizing oxides were yttria (11, 25-26), Calcia (16) and Magnesia (21-24).

The reasons of high fracture toughness due to the presence of intragranular tetragonal ${\rm ZrO}_2$ precipitates are the following :

- 1) Initiation of crack is difficult since work has to be done first to cause the phase transformation and then to initiate the crack.
- 2) The same argument would be true during the

propagation of a crack. As transformation occurs the sudden shape change of the precipitate absorbs strain energy from the crack tip region and a zone of compressive stress is created near the crack tip through which the crack must propagate. Thus, a greater applied stress is necessary for crack growth to continue.

Garvie etal obtained a Ca-psz by a rapid cooling and subsequent ageing treatment. Tetragonal domains which increased in size with time of ageing, were retained at room temperature. The maximum strengthening occured when the tetragonal domains were of a size where they were critically metastable (about 0.1 micron in Ca-psz), i.e. where at room temperature they reverted to a monoclinic form upon the application of mechanical stress.

Similarly, Porter and Heuer have developed 8.1 mole % Mg-psz of optimum mechanical properties, containing homogeneous precipitation of intragranular precipitates. He resorted to special heat treatments of solution annealing of commercial Mg-psz at 1850°C in the cubic SS phase field for 4 hr to dissolve the monoclinic phase present in the as-received material, quenching to less than 1000°C in less than 2 minutes and subsequently ageing between 1400°C and 1500°C, and finally cooling room temperature.

The best materials were those in which the precipitation reaction was nearly two third complete and the precipitates have not lost coherency (size 0.2 mincron in Mg-psz).

 Y_2O_3 psz can be obtained routinely at relatively low temperature (less than 1500°C) (11, 26) and the content of tetragonal phase in the polycrystalline ZrO_2 ranged from 98 to 10% Ypsz has the best properties among all the pszs (2, 11, 27), but Y_2O_3 is a very costly rare earth material.

Hannink (2) has claimed to have produced Capsz, Mgpsz, Ypsz by sintering the 9.98 mole % MgO-ZrO2,8.4 mole % CaO ZrO2 and 9.74 mole % Y2O3 ZrO2 followed by solution treatment in the fluorite phase field at 1800°C, cooling to 1300°C, then quenching to 500°C. After natural cooling from 500°C he aged the samples in air. Ageing temperatures were 1400°C for Mgpsz, and 1300°C for Capsz and Ypsz.

Ageing for a long time (over ageing) causes the tetragonal particles to grow beyond critical size (0.1 micron for Capsz, 0.2 micron for Mgpsz and 0.3 micron for Ypsz depending on the composition). These particles lose coherency with the matrix and transform readily to monoclinic symmetry when cooled.

 ZrO_2 -TiO₂ system has been investigated by several

workers in early 50's but was not thoroughly studied afterwards. The phase diagram by Duwez, Brown and Odell (9) (fig. 2d) shows that the addition of TiO₂ to ZrO₂ causes a rapid decrease of both liquidus and solidus curves, from 2715 to 1820°C, the latter is the incongruent melting temperature of ZrTiO₄.

The addition of TiO₂ also lowers the transformation temperature of zirconia from 1000°C to approximately 350°C. However, stabilization, i.e. formation of cubic phase does not take place.

Voromin (28) has shown that addition of 1% TiO_2 , to stabilized zirconia increases the compressive strength from 1050 Kg/cm^2 to 1450 Kg/cm^2 . Further increase to 2% decreases the strength to 500 Kg/cm^2 . Further addition of TiO_2 increases the compressive strength and at 5% the strength is about as original.

By going to explore the feasibility of making inexpensive electrolytes using the tape process, Radford & Bartton (1976)(6) has added 5 mole % TiO₂ to commercially stabilized zirconia (5 wt % Ca sz & 12 wt % Y₂O₃ sz) so that the tape along with the electrode material, platinum could be sintered to impervious, high density (93%) electrolyte by sintering at 1500°C.

Coughanaur (29) has observed the existence of a single phase solid solution in the $\rm ZrO_2$ rich corner of the $\rm CaO\text{-}TiO_2\text{-}ZrO_2$ phase diagram (fig. 2.e).

The ionic radii of Ca^{2+} , Mg^{2+} , Y^{3+} , Ti^{4+} and Zr^{4+} are shown below (30).

Species	Ca ⁺²	Mg ⁺²	y ⁺³	zr^{+4}	Fe ⁺³	\mathtt{Ti}^{+4}
Atomic number	20	12	39	40	26	70
Atomic radii	1.969	1.594	1.79	1.58	1.241	1.458
Ionic radii (30)	1.06	0.78	1.06	0.87	0.67	0.64
Crystal structure	FCC	Hex		Нср	Bcc	Нср
Ionic radii, (Pauling)0.99	0.65	0.93	0.80		0.68

The average ionic radius of Ca^{2+} & Ti^{4+} is 0.85 aungstrom, close to that of Zr^{4+} . So it is likely to form a solid solution of $Ca0-Ti0_2-Zr0_2$.

It seems probable that compositions containing smaller percentages of CaO than is required to fully stabilize the ${\rm ZrO}_2$ and some amount of ${\rm TiO}_2$ would produce a psz which can retain tetragonal precipitates due to lower tetragonal monoclinic inversion temperature and would have a high fracture toughness, strength and thermal shock resistance.

With the problem of low thermal shock resistance of the glass fiber drawing bushings in view and the above

intension the present study was aimed at improving these thermal and mechanical properties without resorting to any quenching operation.

CHAPTER-II

OBJECTIVES OF INVESTIGATION

- (1) To develop partially stabilized zirconia (psz) of high strength, corrosion resistance and thermal shock (spalling) resistance by addition of (CaO+TiO2).
- (2) To find optimum composition for CaO.psz.
- (3) To study whether TiO2 stabilizes ZrO2 or not.
- (4) To study the stabilization & properties of stabilised zirconia from indigenous monoclinic ZrO2.
- (5) X-ray diffraction studies on the stabilized & partially stabilized zirconia to identify the different phases present (estimate their percentages).
- (6) Comparison of thermal shock resistance of the stabz & pszs.
- (7) To study the corrosion by E-glass on the stabz, pszs ϵ : Al₂0₃.
- (8) Measurement of modulus of rupture in bending for the psz and stab Z's. 22 100 22 100.
- (9) Measurement of hardness and stress intensity factor of the pszs and stabzs.

- (10) Determination of apparent porosity, true porosity and bulk density of slip cast crucibles and pressed samples, Determination of true density, true porosity and closed porosity of pressed samples.
 - (11) Optical microscope studies on stabz and pszs samples after thermal shock resistance test.
 - (12) X-ray diffraction studies to estimate the changes in phase composition during thermal shock resistance test.

CHAPTER-III

EXPERIMENTAL PROCEDURE

3.1 Raw Materials

Monoclinic Zirconia from Indian Rare Earths Ltd., and CaCO3, TiO2, MgO of Laboratory reagent grade were used. For comparison, imported commercial stabilized zirconia, ZircoaB, manufactured by Corhart Refractories Co., U.S.A. was also used to make crucibles and plates.

3.2 Sample preparation

3.2.1 Weighing & Mixing:

Monoclinic zirconia was weighed on a pan balance. CaCo₃, TiO₂& MgO were weighed in a chemical balance. The different batches with different additives were dry-mixed in a ball mill for 2 hrs. for each batch. The compositions are shown in table 3.1.

3.2.2 Precalcination:

The batches were mixed with water and some irregular shapes were made by hand. The batches were precalcined at 1200-1300°C in order to

- a) remove CO2 from the batches carrying CaCO3,
- b) reduce size, c) facilitate handling.

TABLE 3.1

BATCH COMPOSITIONS

Batch	Zro ₂	CaO ·	TiO ₂	Mg O
	mole %	mole %	mole %	mole %
C7	93	7	ét s	43
C10	90	10	6,5	45.8
C13	87	13	«IA	mia
C15	85	15	ca	4ius
C7T5	88	7	5	cus
C8T7	87	8	7	tias (
т7	93	àlige	7	#SAMP
Stab Z (Commercial Zircoa B)	Unknown	rasi	m.s	دے
M8	92	rion.	APCO	8

3.2.3 Calcination

The batches were calcined in a zirconia lined furnace (fig. 3.3) by Indane-Oxygen gas firing. The temperature was raised to 1850°C in 5-6 hrs, & then the temperature was maintained for 4 hrs. The M8 batch was calcined at 1350°C in a globar furnace.

3.2.4 Crushing

primary crushing of calcined lumps were done in the jaw crusher, and then secondary crushing was done in the roll crusher. The particles were reduced to a maximum size of 2 mm. The iron rust particles which came in during crushing were sorted out by in a jnet.

3.2.5 Grinding

Grinding was done in a high alumina porcelain lined ball mill with alumina balls using water as the liquid. Some batches were ground in a agate centrifugal ball mill (Pulverizetle, Fritsch, Germany, Type 05.102) also, The grinding that has been done for different batches is shown in Table 3.2.

The C7T5 and C8T7 batches were ground for a longer time since these developed pour casting properties i.e., quick settling property after 82 hrs of grinding.

TABLE 3.2
GRINDING HOURS

Batch		Hours of grinding			
	in Alumina ball mill	Agote Centrifugal ball mill	Chart speed		
C10	90	umakumakunga Makalah, perintat Perintatangga di kabupatén di penggalapatah ping di Pendah di Pendah di Pendah di Malah	ACC		
C13	82	-			
C7	37	35	5		
C15	may.	50	6		
C71'5	106	-			
C8T7	106				
T 7	18				
м8	20				

The batch T7 was weak and crumbly, so a shorter period of grinding was given to the batch. The M8 batch was calcined at 1350°C and so it was ground for a short time.

3.2.6 Sieving and Magnetic Separation

The slip produced on grinding were passed through a 325 mesh ASTM sieve. Some of the batches were dried and granulated. Then magnetic separation was tried for in a of magnetic separator but the efficiency/separation was poor.

Finally magnetic separation was done by keeping two Alimco bar magnets in the slip kept in a plastic bucket and stirring the slip by a 3 blade stirrer. The magnetic particles adhering to the edges and corners of the magnets were wiped out from time to time at 10-15 minutes interval by cotton. This process was continued till insiginficantly small amount of magnetic particles got adhered to the magnet over a long time interval.

3.2.7 Preparation of Plaster of Paris Moulds

A wooden model was cleaned and brushed with soap solution. Then it was oiled with coconut oil by a brush. The model was enclosed in a thick sheet of paper folded into a cyclindrical form and was bound with threads. The plaster of paris slip was poured on to the model and it was allowed to set for 30 minutes after which the mould was released from the model. Several moulds (fig. 3.4) were made in this way.

3.2.8 Slip Casting

Slip casting was done for all the batches. The C8T7 batch settled very quickly and a small amount of $\mathrm{Na_2^{CO}_3}$ (~ 0.1%) was added before slip casting whereby the casting properties improved. The T7 batch also settled very quickly

but casting was done without any addition.

3.2.9 Pressing

Pressing was done in a hydraulic press with a pressure of $14.2 \, \text{KN/cm}^2$ (20610 psi).

The powder was mixed throughly with 2% PVA solution in a quantity to have proper pressing characteristics, i.e., easy mould release, proper compaction, avoidance of extra water release during pressing and avoiding crumbliness of the pressed sample. The moist powder was put in the high chromium high carbon alloy steel mould (fig. 3.1) - having an inner dimension of 5 cm x 1 cm x 1 cm. It was levelled by the punch. Then the mould with the punch was kept on the platform of the hydraulic press. Pressure was applied at a slow rate (in ~ 2 minutes). After attainment of the maximum load, 2 minutes were allowed before the release of the pressure in order to homogenize the pressure distribution. The sample was released in a hydraulic hand press.

The percentage of water added was determined on the dry basis and are shown in the table.3.3 below.

Table 3.3 Water of Granulation for Pressing

Batch	Water % on dry basis	Batch Water	% on dry basis
C7T5	9-11	Clo Clo	4.3-4.5
C7	6.4-7	Т7	6.7-7.3
C10	4.9-5	M8	3.2-4

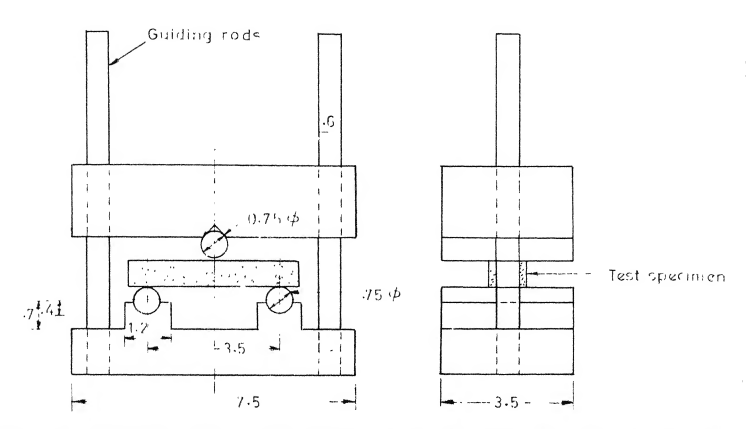


FIG. 3.2 FRONT VIEW AND SIDE VIEW OF THE BEARING EDGES FOR MODULUS OF RUPTURE TEST

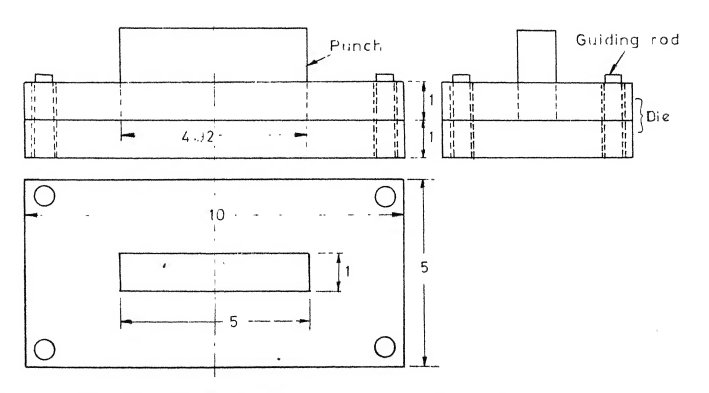


FIG. 3-1 DESIGN OF THE MOULD ASSEMBLY USED FOR PRESSING.

FIG.33 PLASTER OF PARIS MOLDS USED FOR SLIP CASTING

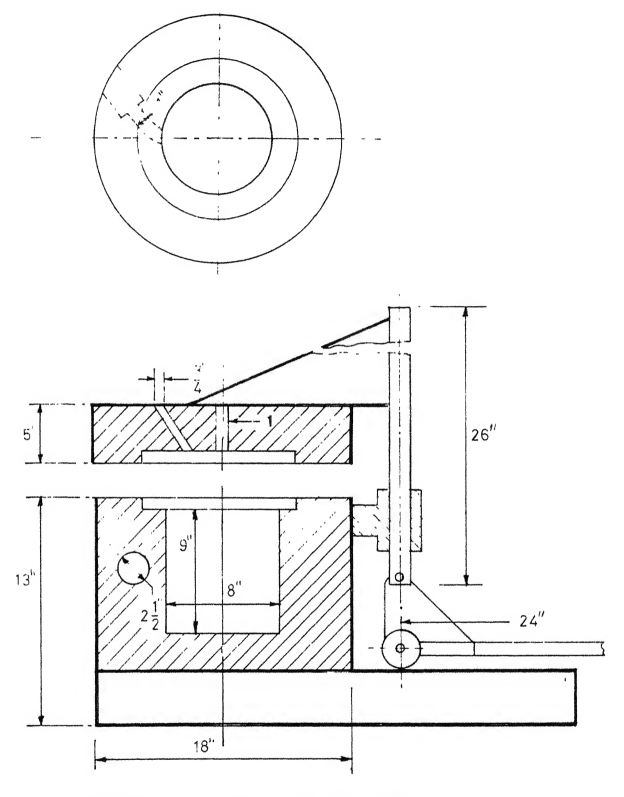


FIG. 3.4 ZIRCONIA LINED FURNACE

3.2.10 Drying

The slip cast crucible were air dried and the pressed samples were dried in an oven at 180°C.

3.2.11 Presintering and finishing

The crucibles were presintered at 800°C in an annealing furnace for 5 hrs. The rims were polished by rubbing on a plain paper.

3.2.12 Sintering

The test bars and crucibles were set in the zirconia lined furnace (fig. 3.3). Indane-oxygen gas firing was used. Temperature was raised to 1950°C in 14 hours and soaking of 1 hr was given at that temperature. The temperature was reduced to 1350°C in 12 hrs and then the gas firing was shut off, and the furnace allowed to cool by itself.

3.3 X-ray Diffraction Studies

Solid bars of the sintered samples and samples after thermal shock resistance test were mounted on the X-ray diffractometer. The operational variables are listed in table 3.4. The angles at which well defined peaks appeared were noted down and the corresponding 'd' values were calculated from the Bragg equation,

$$n_{\lambda} = 2d \sin \theta$$
 (3.1)

TABLE 3.4

PARAMETERS USED IN X-RAY DIFFRACTOMETER

Radiation used : CuK, \sim . ($^{\circ}$ = 1.5405 Å)

Filter : Ni

Voltage rating : 35 KV

Current : 12 mA

Intensity range : 1000 counts per second

Time constant : 2 seconds

Beam slit : 3°

Detector slit : 0.2°

Scan speed : 2° per minute

Chart Speed : 2° per minute

Speed controlling disc : C

where n = an integer, taken as 1. = wave length of X-ray radiation, $\Theta = Bragg$ angle, d = inter planar spacing.

The d'values & the relative intensities from peak heights were compored with the standard powder diffraction file (PDF) in order to identify the crystalline phases. the integrated area of the 100I peaks of different phases were determined by trapezium method by using the following equation (3.2),

Area, A = h.
$$\frac{f_1}{2} + f_2 + f_3 + f_4 + \dots + \frac{f_{n+1}}{2} \dots (3.2)$$

where h = length of one division

n = number of divisions

and f_1 , f_2 , f_3 , f_{n+1} are the respective functions at the divisions, 0, 1, 2, n.

The phase analysis was done according to the modified formula in polymorphsmethod developed by Carvie & Nicholson Volume fraction of monoclinic phase,

$$V_{m} = \frac{1.603 \left[I(11\overline{1})m\right]}{1.603 \left[I(11\overline{1})m + I(111)c\right]} \dots (3.3)$$

where I $(11\overline{1})_{m}$ = integrated intensity of the $(11\overline{1})_{m}$ monoclinic reflection.

 $I(111)_{C}$ = integrated intensity of the (111) cubic reflection and it is assumed that

$$V_{m} + V_{C} = 1$$
(3.4)

 V_{C} = Volume fraction of cubic phase

 V_{m} = Volume fraction of monoclinic phase.

3.4 Thermal Shock Resistance (Spalling resistance) Test

A globar furnace with thick Sigr rods as heating elements and high duty fireclay lining was used for the test. The furnace can regain the temperature drop during removal of the samples in about 10 minutes. The rectangular bar test pieces were kept in a refractory support and placed inside the furnace. The furnace was heated at a uniform rate so that 1300°C was attained in 4 hrs. The temperature was maintained for 30 minutes and then the support containing the samples was removed from the furnace with large tongs. The samples were placed on a brick in a position free from draughts. After they had been cooled for 10 minutes they were examined and then replaced into the furnace for a further period of 10 minutes. Then the cycle was repeated. test was concluded when the specimens could be pulled apart by light tongs. One highduty insulation brick support could be used for two cycles.

3.5 Corrosion Resistance Test

The E glass marbles were melted in a sillimanite crucible and quenched into water to get small particles of

the glass. The crucibles were filled with the E glass and put on the grooves made by drilling in a highduty insulation brick support. The support with the crucibles was kept in the globar furnace and heated to 1400°C in 8 hrs. The temperature was maintained for 13 hrs. Then the furnace was cooled down to room temperature. The bottoms of the crucibles were cut into two vertical cross sections by diamond tipped wheel. The two sections were broken apart by a small hammer. Photographs of the cross sections of the crucibles were taken and corrosion of the crucibles were studied.

3.6 Determination of apparent porosity, apparent (bulk) density, true porosity & true density. (32)

3.6.1 Apparent porosity & bulk density.

The test specimens were dried at $110\,^{\circ}\text{C}$ and weighed (") after cooling to room temperature in a desiccator. The test specimens were placed in distilled water and boiled for about 2 hrs. then these were allowed to cool to room temperature, while still immersed in water. The test specimen was weighed while suspended in water (W_2). Immediately after obtaining the suspended mass, the test specimen was removed from water, blotted lighly with a wet tissue paper and weighed in air (W_1).

Then apparent porosity & apparent density were calculated from the following formulae:

Apparent (bulk) density,
$$D_A = \frac{W}{W_1 - W_2}$$
(3.5)

Apparent porosity,
$$P_A = \frac{W_1 - W}{W_1 - W_2} \times 100$$
(3.6)

3.6.2 True Specific gravity, true density and porosity

3.6.2.1 Sample Preparation

Samples weighing about 20 gms were ground in an to such a fineness that they would pass through 149-micron IS-Sieve. The sample was dried at 105 to 110°C.

3.6.2.2 The pronometer was washed, dried and weighed (P). About 4 to 6 gms of sample was placed in the dry pronometer and weight of pronometer, stopper plus sample (W) was taken.

The precommeter was filled to one-half of its capacity with distilled water which had been boiled to remove dissolved air and cooled. The pycnometer was kept on a hot plate to boil the water for 10 to 15 minutes. Then the pycnometer was kept in a distilled water bath in a glass petri-dish to cool the pycnometer to room temperature. The pychometer was filled with water, the stopper was inserted and the excess water was wiped off from the stopper. The

pycnometer was throughly dried with a tissue paper. Then pychometer & the contents were weighed (W₂). The sample was thrown away, the pycnometer was washed and filled with water, and it was weighed after drying the outside with tissue paper.

The specific gravity, true density and true porosity were calculated from the following formulae :

Specific gravity,
$$S = \frac{W - P}{(W-P) - (W_2 - W_1)}$$
(3.7)

where

W = mass in gm of the stoppered pycnometer and sample.

P = mass in gm of the stoppered pychometer

 W_2 = mass in gm of the stoppered pychometer, sample and water.

 W_1 = mass in gm. of the stoppered pycnometer filled with water.

True density = $Sp. gr. x (d_w - d_a)$

where $d_{\rm W}$ = density of water at the temperature at which the test was carried out, and

d_a = density of air at the temperature at which the
 test was made.

True porosity,
$$P_{T} = 1 - \frac{Apparent density, D_{A}}{True density, D_{T}}$$
(3.8)

True porrosity, percent =
$$(1 - \frac{D_A}{D_T}) \times 100$$
(3.3)(32)
Closed porosity P_C (percent) = P_T (%) - P_A (%)(3.10)

3.7 Microstructural Studies by Optical Microscopy

3.7.1 Sample Preparation

The samples were joined together into two batches by quick fix in order to polish a few samples together. These were polished with Sic Powders of mesh sizes 220, 400, 600 and 800 successively. After this optical polishing was done on a polishing wheel with diamond paste. Little amount of liquid paraffin was added to the diamond paste and also added again for 2-3 times during polishing. Final polishing was done with 0.3 micron and 0.05 micron Al_2o_3 powders successively to remove the pits in the sample produced during polishing by diamond paste.

3.7.2 Microstructure

Photographs were taken with the microscope focussed at a magnification of 200X and 400X for sintered samples and 400X for the spalled samples. Camera magnification was 1/3rd the microscope magnification. An exposure time of 3 seconds was given in each case. The photographs after enlargement during printing were studied for different phases, grain structure and changes on spalling (thermal shock).

3.8 Particle size analysis of the ground batches

Different solvents were tried to disperse the particles and a mixture of Ethyl alcohol, methyl alcohol and acetone was found to be the best. The mixed solvent was taken in an agate mortar in small amount. A pinch of the powder was added to it. It was mixed with the liquid by whirling the paste. Then the mortar with the contents was placed in the water bath of an ultrasonic cleaner and the bath was vibrated with ultrasonic waves for ten minutes. Immediately after this one or two drops of suspension were dropped on a cleaned glass slide with the help of a dropper and allowed to spread and dry in air.

The slide was kept in a petri dish. The photograph of the particles were taken with 400 X magnification in the microscope, combined with the 3 times reduction by the camera. Printing of the photographs was done with an enlargement of 3.5. Each photograph of different batches was divided into several areas and the size of the particles of each area was read by a scale. The number of particles in different size ranges were obtained and particle size distribution was thereby found out.

3.9 Modulus of rupture

The sintered bars were ground with 220 mesh SiC powder to give proper rectangular shape and an approximate dimension of $4-4.5 \times .9 \times .7 \text{ cm}$. The samples were polished with SiC powders of mesh sizes 400, 600 and 800 successively.

The samples were tested in the Instron machine with the bearing Edges for modulus of rupture test, shown in figure (3.2).

The load cell used had a range of 0-200 Kgs with a least count of 2 Kg. The cross head speed used was 0.2 mm/min. Chart speeds of 0 cm, 2-0 cm, 5 cm per minute were used.

The Instron was first calibrated to read 200 Kg in the full span of the chart.

The sample was mounted on the bearing edges of the Modulus of rupture test jig (fig. 3.2) kept on the platform of the Instron under the cross head.

The load was applied by downward movement of the cross head. The maximum atbreaking load after which the specimen failed suddenly was obtained from the chart.

From the breaking load, the modulus of rupture was calculated from the following formula :-

Modulus of Rupture, in
$$Kgf/cm^2 = \frac{3Wl}{2bd^2}$$
(3.11)

where, h = load in Kgf at which the specimen failed,

l = distance in cm between the centre lines of the lower bearing edges,

b = width of the specimen in cm, and

d = depth of specimen in cm.

3.10. Vickers Hardness and Fracture toughness

The hardness was measured by indenting polished samples using a square base diamond pyramid as indentator (33) in the vickers pyramid herdness testing machine (Vickers Instruments Ltd., England, model No. 255038).

Because of the shape of the indenter this is frequently called the diamond pyramid hardness test. The diamond pyramid hardness (DPH) or Vickers hardness number (VHN or DPH) is defined as the load divided by the surface area of the incentation.

$$IPH = \frac{2P \sin(Q/2)}{I^2} = \frac{1.854 P}{L^2} \dots (3.12)$$

where P = applied load, Kg.

L = average length of the diagonals, mm.

 Θ = angle between opposite faces of diamond = 136°.

The fracture toughness was determined by a technique developed by Evans & Charles (34) using the dimensional analysis of indentation crack length (c) and impression

radius (a).

Fracture toughness, k is obtained from the following formula :-

$$k_c \phi/H \sqrt{a} = 0.15 k (\frac{c}{a})^{-3/2}$$
 ...(3.13)

where k = fracture toughness

 ϕ = the constraint factor (\sim 3)

H = hardness

a = impression radius

c = crack length

k = a correction factor

The function $\frac{k_c}{H\sqrt{a}}\left[\frac{H}{E}\right]^{0.4}$ when plotted with $\frac{c}{a}$, produces a unique curve which fits for all the polycrystalline materials. The straightline curve for single crystals fall close to those of polycrystals.

The samples which were used for studying the microstructure were taken. Opposite surface to the polished one was ground in 220 mesh SiC powder to make the two surfaces parallel. In the vickers hardness testing machine loads of 2.5 kg and 5 kg and were used for the test. For each load several indentations were made. In each case the diagonal of the indentation, the average crack length of the cracks produced by indentation were measured by ocular reading, one coular reading corresponds to

0.001 ma. .

Hardness corresponding to each reading was read off the vickers chart. Hardness values in ${\rm Nm}^{-2}$ unit was also calculated from equation (3.9) for use in determining ${\rm k}_{\rm c}$.

Using the measured impression radius, cracklength and hardness, the fracture toughness was determined from the calibration curve (34, 13).

Photographs of some of the indentations were taken.

CHAPTER-IV

RESULTS AND DISCUSSION

4.0 Particle Size analysis

The results of the particle size analysis of the ground ZrO_2 powder before shaping are shown in table 4.1. The particle size distributions are shown in the figures 4.1, 4.2, and 4.3.

All the batches have highest distribution density at less than or equal to 0.214 micron size. The maximum particle size observed is 3.65 micron. So the powders are very fine. This is because of the fact that the batches have been ground for a very long time in order to develop good casting properties.

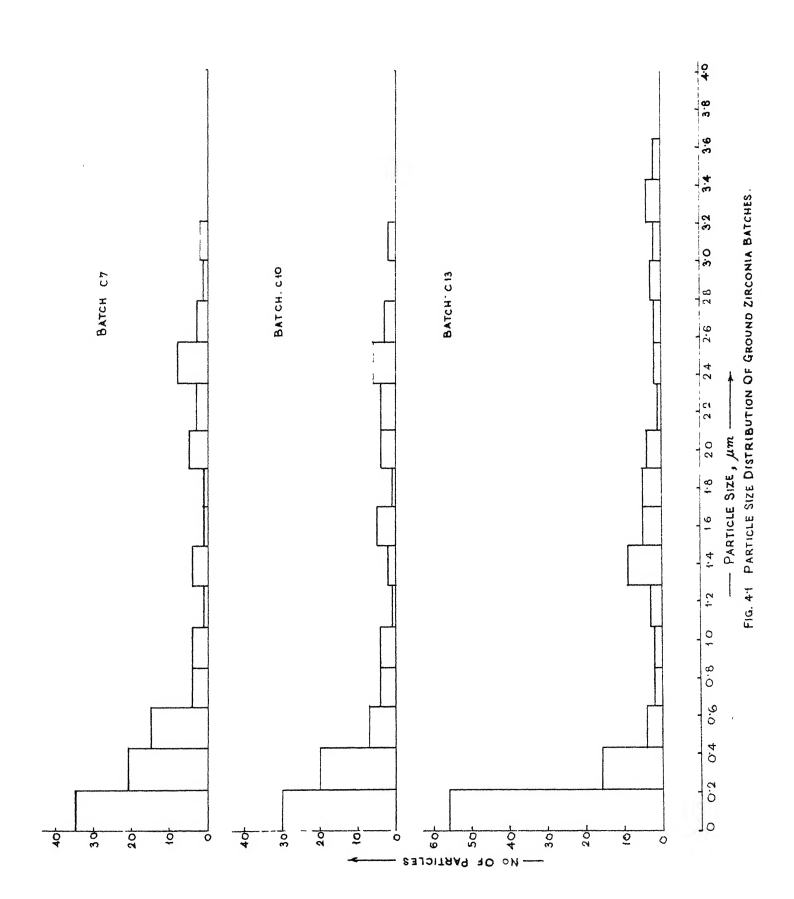
4.1 Densities and porosities

The bulk density, true density, apparent porosity, true porosity and closed porosity values of the different batches are recorded in table 4.2, and 4.3. The variation of true density and true porosity values as a function of composition are shown in figures 4.4 and 4.5 respectively for sintered bars made by pressing and figures 4.6, 4.7 for crucibles made by casting. The bulk density is much less than the true density and most of the porosity is due to

TABLE 4.1

PARTICLE SIZE ANALYSIS OF THE GROUND BATCHES

Size range, micron	No.	of I	Particle		ling i atches		size	range	a Tapatah berbail dan punda da d
ланирадзініцияння намен таком у чт і зако жикладані правинанция	С7	C10	C13	C15	C7T5	C8T7	Т7	Stabz	M8
₹0.214	35	30	56	19	43	32	73	63	126
0.214- 0.429	21	20	16	23	25	27	28	57	96
0.429- 0.643	15	7	4	16	17	13	17	29	53
0.643 0.857	4	4	2	3	16	3	27	11	15
0.857. 1.072	4	Ą.	3	4	3	4	26	7	9
1.072- 1.286	1	1	3	2	1	4	20	9	7
1.286- 1.499	3	2	9	د ه	3	2	11	6	4
1.499 1.714	5	5	7	1	1	4	18	5	3
1.714- 1.929	1	1	5	2	1	• •	11	4	1
1.929- 2.143	5	4	4	6	2	2	19	11	2
2.143- 2.357	3	4	1	5	2	4 ==		5	فندة
2.357- 2.571	8	6	2	9	1	1	11	5	۳
2.571- 2.786	3	3	2	1	G 3	1	دب	3	
2.786- 2.999	1	£3	3	1	æ a	1	7	1	1
2.999- 3.214	2	2	2	2	د. ع	engos	1	دسه	وستنه
3.214- 3.429	rs.	ea	4	دنه	E.3-	Sec. 3	eب	فسة	رى
- 4.286	+43	4	e.a	1	ه. ع	1	دء	E,	



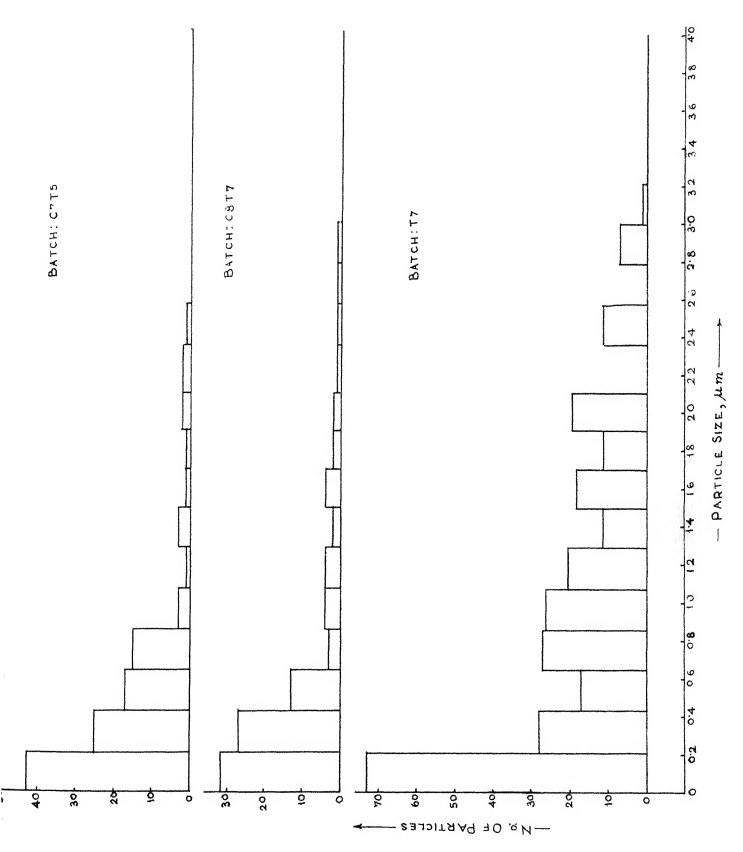


FIG 4.2 PARTICLE SIZE DISTRIBUTION OF GROUND ZIRCONIA BATCHES.

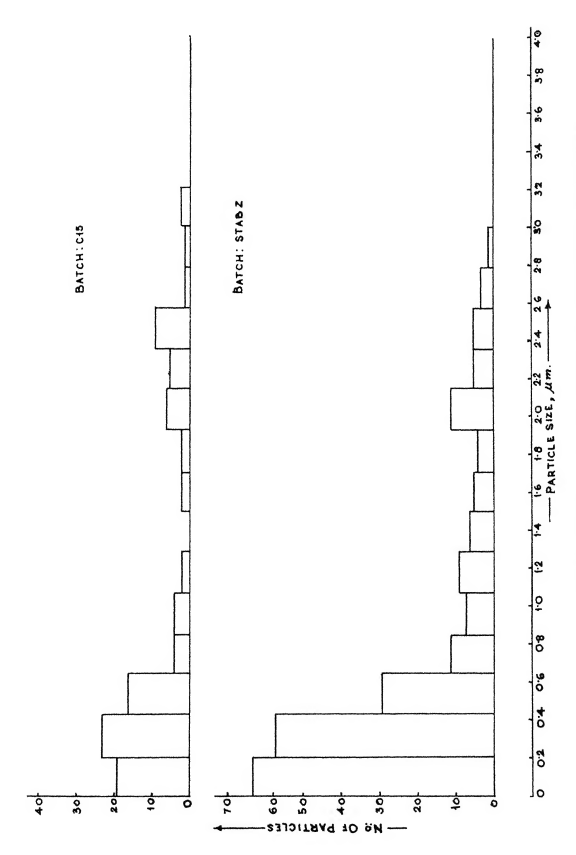


FIG. 4:3 PARTICLE SIZE DISTRIBUTION OF GROUND ZIRCONIA BATCHES.

TABLE 4.2

TRUE DENSITY, BULK DENSITY, APPARENT, CLOSED AND

TRUE POROSITIES OF THE SINTERED SAMPLES

					The same of the sa
Sintered bars	True density D _T . gms/cc	Bulk density D _{B'} , gms/cc	Apparent porosity PA %	Closed porosity PC %	True porosity ^P T %
C7	5.364	4.794	1.076	9.554	10.63
	5.373	5.083	1.910	3.429	5.399
C10				3.404	5.006
C13	5.404.	5.133	1.620		4.115
C15	5.572	5,300	0.000	4.115	
C7T5	5.475	4.730	1.457	12.153	13.610
C8T7	5.541	4.82 2	1.603	11.377	12.980
T7	5.486	4.733	1.020	12.703	13.723
M 8	5.246	4.320	3.730	13.921	17.651

TABLE 4.3

APPARENT DENSITY, BULK DENSITY, APPARENT CLOSED

AND TRUE POROSITIES OF THE CRUCIBLES

Crucibles	Bulk density ^D B gms/cc	Apparent porosity P _A %	True porosity ^P T %	Closed porosity ^P C %
and the second second section of the second section of the second second second section sectio	anadalaidentajaudekaggydickon a modujadek Trenia dekolektrajaisieki. He	aker paraget		
C7	4.914	6:946	8.380	1.434
C10	5,038	1.665	6.237	4.572
C13	5.153	1.715	4.631	2.916
C15	5.370	0.000	3.625	3.625
C7T5	4.882	2.785	10.830	8.045
C8T7	5.101	2.126	7.930	5.810
т7	4.998	3.354	8.892	5.538
M8	4.642	7.300	11.514	7.300
Stab z	5.062	2.926	8.562	5.636

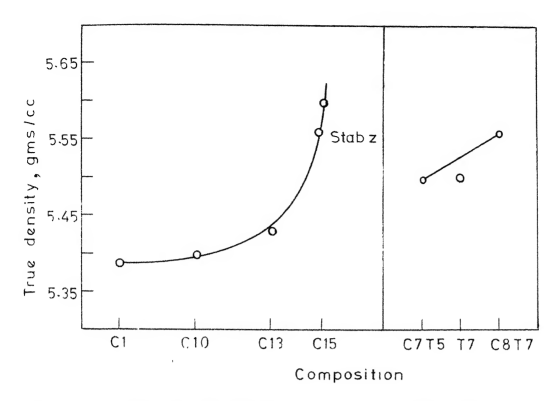


FIG. 4.4 TRUE DENSITY VS. COMPOSITION

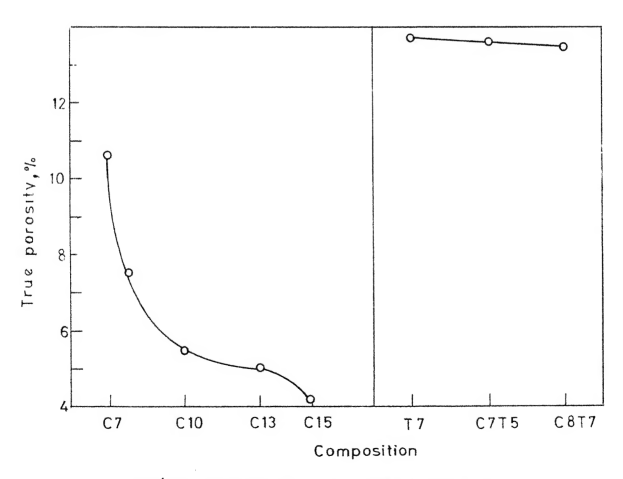


FIG. 4-5 TRUE POROSITY VS. COMPOSITION

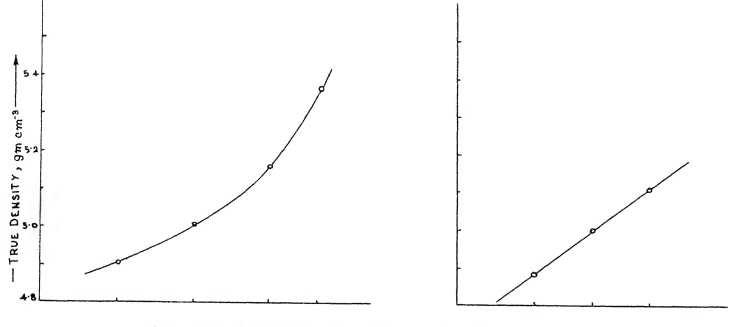


FIG 46 TRUE DENSITY VS. COMPOSITION FOR CRUCIBLES.

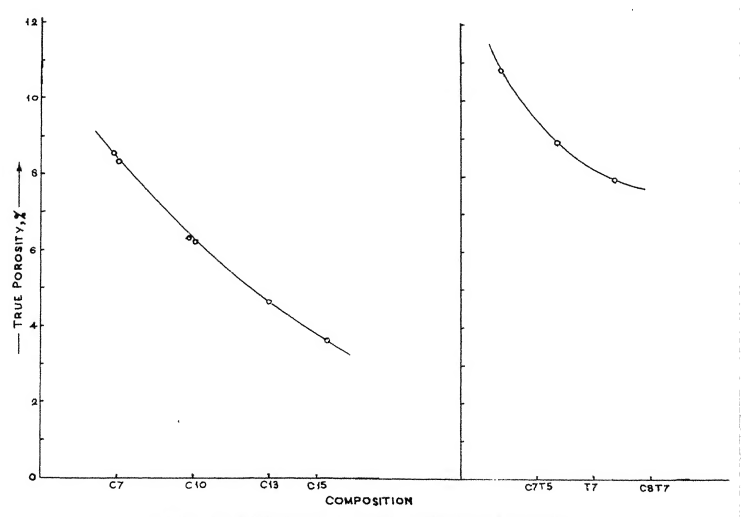


FIG. 47 TRUE POROSITY VS. COMPOSITION FOR CRUCIBLES.

closed pores as true porosity is much greater than apparent porosity.

The true porosity ranged between 4.1 for C15 to 13.7 for C7. The M8 batch has a very high porosity 17.65 since it was calcined at 1350°C and on sintering, the samples swelled to some extent.

The porosity figures of CaO-TiO₂-ZrO₂ and TiO₂-ZrO₂ batches are much higher than the CaO-ZrO₂ compositions. The reason is as follows. Titanium has 4+ valency. So when it will replace Zr⁴⁺, no vacancy will be created as opposed to Ca²⁺. In the latter case vacancies are formed to maintain charge balance. So in case of Ti⁴⁺ substitution diffusion will be lower than that in the case of Ca²⁺ substitution due to lower number of vacancies in CaO-TiO₂-ZrO₂ composition.

Density increased with increasing amount of CaO as more and more liquid phase sintering occured and also at the same time number of vacancies is increased.

Porosity of the crucibles is higher than porosity of the sintered bars because the crucibles were made by slip casting whereby the green density was less than that in the case of pressed bars.

4.2 X-ray Diffraction Studies

The measured d'values, respective intensities of the

different lines, integrated intensity and phases present in the sintered samples are shown in tables 4.4a to 4.4i. The X-ray data of the batches after thermal cycling (spalled) are recorded in the tables 4.5a-4.5i. The X-ray diffraction curves of sintered and spalled specimens are shown in figure 4.4(i)-(iii) and 4.5 (i)-(iii) respectively.

By comparing the measured 'd' spacings and ${\rm I/I}_{\rm O}$ values of our samples with those of ASTM standard we have come to the following conclusion.

The C15 batch contained 95.73% cubic phase and the rest as monoclinic phase, that is the stabilization in this composition was almost complete. In case of C13 batch the cubic phase was only 30.73% indicating the importance of the CaO amount for stabilization. The C10 composition did not get stabilized to the cubic phase. It contained 15.5% tetragonal and 84.5% monoclinic phases respectively.

The C7 composition had only a small amount of tetragonal phase (3.06%) and monoclinic (96.94%) was the major phase.

Lower amount of CaO resulted in higher amount of monoclinic form. T7 composition had produced a monoclinic solid solution as indicated by slight change in'd' values for all (hkl) planes. This agrees with the phase diagram, figure, 2.d.

C7T5 batch also formed a single phase monoclinic solid

solution which agrees with the phase diagram, figure 2e.

C8T7 composition had developed a small amount of tetragonal phase (5.8%). The M8 batch was also fully monoclinic.

On repeated thermal cycling more and more phase had become ill defined and showed broad K-ray peak and some of them formed humps instead of well defined peaks. This means that the 'd' spacing has been changed, crystallinity has got reduced and micro cracks have been produced throughout the matrix. The C7T5 composition retained some tetragonal phase. It seems that this composition has a low temperature monoclinic-tetragonal phase boundary. The investigation of the phase diagram is necessary in this system to clarify this point.

The composition C7, C10, C13, C15 and Stab Z had also retained tetragonal phase. But retention of tetragonal phase was highest in the two compositions-C7 and C7T5.

On thermal cycling the diffracted intensity from some of the hkl planes increased and the intensity of some of the planes decreased.

of

The decrease of amount/cubic phase and formation of monoclinic phase is possibly because of the following reason.

At the furnace temperature of thermal shock test the .
C15 and Stab Z samples got heat treated in the (cubic+monoclinic)

TABLE 4.1
ASTM X-RAY DATA FOR CUBIC, TETRAGONAL AND MONOCLINIC ZIRCONLA

Cubic, a	o = 5.03		Tetragonal	$a_0 = 5.07$,
		_		c _o = 5.16
d Å	I/I _o		d A	I/I _o
2.96	100		2.93	100
2.56	24		2.52	AO
1.81	80		1.81	60
1.54	60		1.79	100
1.48	10		1.55	50
1.28	12		1.53	100
1.17	20		1.47	60
1.04	16		1.29	40
0.98	14		1.27	60
0.90	5		1.17	40
0.86	8		1.13	40
0.85	5		1.10	60
0.81	4		1.05	40
			1.04	70
			0.99	40
			0.98	70

Monoclinic

		Monoclinic			
೧ =	5.1477,		b _o = 5	5.2030,	
c _o =	5.3156			99°23 °	
d A	I/I _o		å Å b	I/I _o	-
5.036	6		1.656	14	-
3.690	18		1.640	8	
3.630	14		1.608	8	
3.157	100		1.591	4].	
2.834	65		1.581	41	
2.617	20		1,541	10	
2.598	12		1.508	6	
2.538	14		1.495	10	
2.488	4		1.176	6	
2.328	6		1.447	- تاریخ	
2.285	2		1.420	6	
2.252	4		1.358	2	
2.213	14		1.348	2	
2.182	6		1.321	6	
2.015	8		1.309	2	
1.989	8		1.298	2	
1.845	18		1.269	2	
1.818	12		1.261	2	
1.801	12				
1.780	. 6				

14

1.691

TABLE 4.4

X-RAY DIFFRACTION DATAS OF SINTERED SAMPLES

c = cubic, m = monoclinic, t = tetragonal.

Table 4.4 a. -- Composition C15

		•	· ·		· · ,
20 degrees	d A	I I _O	Integrated Intensity	Remarks	% phases
27.2	3.2756	6.0	600 VIII.	_{ಲಾ} ಜು	ھي جب
28.2	3.1617	2.25	4.40	m	4.271
30.26	2.9511	100	158.085	С	95.729
35.08	2.5558	15.02	est) ette	С	Kana (120
50.33	1.810	39.04	8678 hand	С	2000
59.90	1.5428	18.92	equipa t. 114	С	حنيها فتكله
62.72	1.4801	5.105	eza e >	С	es is

Table 4.4.b. -- Composition C13

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% phases
24.2	3.6746	14.35	Las r -	m	ಬ್ವ
24.55	3.6230	14.57	con 129	m	Cath Co
28.34	3.1464	100.00	135.98	m	69.27
30.24	2.9530	60.87	96.7	C	30.73
31.58	2.8307	80.44	ब ्च देश	m	Aciy Kiri
34.40	2.6048	32.61	628 C-1	m	فيجسا
35.40	2,5334	30.44	فسنة وفقات	m,C	Ca tie
38.63	2.3287	7.17	مناوع	m,	ACM CTO
40.88	2,2055	21.74	ಭಾಷು	m	septe etc.a
45.02	2.0119	11.96	The book	m	£255 &2-3
45.62	1.9868	13.48	ning april	m	هنمه مشت
49.52	1.8391	33.70	eus c.J	m	gra 4a
50.30	1.8124	65.22	ಕ್ಷಮಾ ಕಿ.ಮೇ	m	e, sera
54.16	1.6921	18.26	600 G.J	m	هنته هنته
55.57	1.6521	23.91	భ్రా డు	m	هيريه حريبه
60.00	1.5405	38.48	SECS ACLA	m,c	espo dist
62.93	1.4757	24.57	acon 4429	m,c	23 CJ
64.30	1.4475	J 8. 26	وين دين	m	and the
65.80	1.4180	21.09	1689 Q.C	m	జాన సూ
69.10	1.3582	4.78	28 CL)	m	
71.10	1.3248	13.91	g238 471-a	m,	egg-eth

Table 4.4.c.-Composition C10

20 degrees	d A	I	Integrated Intensity	Remarks	% Phases
24.32	3.6567	14.82		m	
24.74	3.5960	9.56		m	
25.63	3.4727	3.46		m	
28.43	3.1367	100.00	145.20	m	84.51
30.41	2,9368	20.43	42.65	t	15.49
31.68	2.8220	58.80		m	
34.33	2.6099	30.15		m	
35.49	2.5272	14.33		t,m	
36.10	2.4859	6.92		m	
38.86	2.3154	8.24		m	
41.08	2.1953	21.90		m	
45.12	2.0051	11.50		m	
45.68	1.9843	14.33	1	m	
49.48	1.8405	36 . 57		m	
50.42	1.8084	37.89		t,m	
50.83	1.7945	39.04		t,m	
54.28	1.6885	18.95		m	
55.61	1.6512	44.95		m	
57.41	1.6037	11.04		m	
58.32	1.5808	11.86		m	
60.12	1.5378	28.34		m,t	
				Contd	

Contd....C10

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% Phases
62.14	1.4925	20.59		m	
02.414	1.4923	20.39		111	
62.86	1.4772	16.31		m	
64.60	1.4414	7.91		m	
65.78	1,4184	15.32		m	
71.40	1.3200	14.50		m	
75.40	1.2596	9.89		m	

Table 4.4.d. -- Composition C7

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% phases
24.27	3.6641	13.31	es (s	m	لب د د
24.68	3.6042	8.40	Cas Cas	m	وستا فسا
25,60	3.4767	9.10	ब्याव ६००३	m	جسه جب
28.40	3.1399	100.00	177.52	m	96.936
30€50	2-9283	5.41	8.995	t	3.064
31:63	2.8263	52.80	an en	m	آ. ا دری
3 4 : 29	2.6126	29.40	വഴ	m	دے دے
35.56	2.5214	10.36	೧೩೫ ನ್ 1	t	مستا دشت
36.00	2,4926	7.28	eag F.P	m	Chi di Good
38.86	2.3154	6.30	can mea	m	من س
40.98	2.2004	21.00	فبنك شيته	m	eap +3
45.02	2.0119	9.38	etas vais	m	فد ما البيدة
45.73	1.9823	12.05	etu (s)	m	فسنة جينت
49.54	1.8382	35.01	6-46 3	m	فيبه فبيط
50.42	1.8084	30.80	es kā	t	ens f 1
50.75	1.7974	32.20	نية جي	t	Star Lad
54.10	1.6937	21.00	ఒద గచ	m	لائية خستة
5 5.63	1.6507	47.48	es co	m	23 ()
5 7 , 39	1.6043	11.20	ಮು ಕಗ	m	ध्य ७७
58,46	1.5770	13.30	ttula Alme	m	السنة الشيئة
60.23	1.5352	25.21	M. D. C. A	t	لنسنا هنتا
61,63	1.5037	14.28	6364	m Contd	C7

Contd....C7

20 degrees	đ Å	I I _O	Integrated Intensity	Remarks	% phases
62.23	1.4906	20.73	caca	m	د ، د
63.02	1.4738	13.59	care a	m	دسه التجا
64.57	1.4420	7.98	e-se-a	m	فينة حينة
65.80	1.4180	13.87	مسة فحشه	m	فرسا فسنا
69.37	1.3536	3,92	wes	m	ఓపా కలు
71.37	1.3205	16.81	413 STP	m	emp 4423

	Table	4.4.e Comp	position Stab	Z	
20 degrees	å Å	I	Integrated Intensity	Remarks	% phases
27.42	3 .2 499	3.98	دعدع	m	ما المائة
28.30	3.1508	2.16	3.5	m	3,036
30.26	2.9530	100.00	179.2	С	96.964
35.11	2.5560	14.10	ent a	С	ette e. J
45.43	1.9947	1.99	భావు కడని	m	డు జు
50.22	1.8151	43.12	era exa	m , c	Cap 40s
59.73	1.5468	5.64	इंद्रा क्ष	C	N RELACION
60.22	1.5355	18.24	هنبه	С	فبسا فيباث
62.30	1.4782	4.98	മാ വ	С	فهيئة مخشقة
74 .5 3	1.2721	2.98	e2 e	m,c	KID CO

Table 4.4.f.-- Composition C7T5

20 degrees	d A	I	Integrated Intensity	Remarks	% phases
24.05	3.6971	15.77	دے دے	m	9 40
24.47	3.6314	10.51	دے چھو	m	استهد ا
25.29	3.5168	5.12	SCOR C. >	m	
28.28	3.1529	100.00	ca c J	m	جه د ع
31.48	2.8394	56.74	ജാധ	m	కమి కనిక
34.12	2.6251	35.04	ಬು ಟಿ	m	ಕುಬ
35,30	2.5400	10.78	423 ° is	m	Marie Activ
35.83	2.5040	7.41	ब्य ट्या	m	కు ఈ
38.60	2.3300	8.63	ينين	m	9.00
40.79	2.2154	18.46	خ کا لندی	m	ats 429
44.76	2.0274	11.59	ھيڪ جين	m	فثيه هيكة
45.48	1.9926	15.50	can tau	m	డా బు
49.26	1.8489	36.52	فنري ووون	m	बंद्धा ८५३
50.23	1.8148	32.75	ब्देश शास्त्र	m	Fra.J 6253
50.57	1.8034	36.12	బాబు	m	د- <u>با ج</u> ب
51.20	1.7827	12.40	activ Allah	m	فيته لنسا
54.02	1.6960	15.23	فيبه (ب	rn	جيئ دي
55.41	1.6567	36.79	ente cus	m	8273 Yuu3
57.24	1.6081	11.05	Adap CI	m	تنه سه
58.18	1.5843	15.23	ಒದ ಮಾ	m	8139 L-79

Contd....

Contd....

Contd....C7T5

29 degrees	d Å	I I _O	Integrated Intensity	Remarks	% phases
59.79	1.5454	29.65	City full	m	andersandersker der der der der der der der der der d
61.35	1.5098	14.83	May sus	m	د)
61.93	1.4970	20.22	High C.Es	m	هن س
62.80	1.4784	11.73	表演 にご	m	రావు ఇద్దు
64.13	1.4509	5.39	ED . 1	m	دے دے
65.63	1.4213	15.23	డుకు శంచి	m	ھن ھن
69,00	1.3599	3.77	cits ray	m	ದಾ ಚಾ
71.18	1.3235	13.75	دے والے	m	(23 63)
20 degrees	d Å	g Com	oosition C8T7 Integrated Intensity	Remarks	% phases
20 degrees	0		Integrated	Remarks m	% phases
Secretario de la constitución de	d Å	I I _O	Integrated Intensity		
24.04	d Å 3.6986	I I ₀	Integrated Intensity	m	
24.04 24.63	3.6986 3.6115	18.08 13.56	Integrated Intensity	m m	ca co
24.04 24.63 25.44	d Å 3.6986 3.6115 3.4982	18.08 13.56 5.79	Integrated Intensity	m m m	E3 E3
24.04 24.63 25.44 28.20	3.6986 3.6115 3.4982 3.1617	18.08 13.56 5.79 100.00	Integrated Intensity	m m m	94.215
24.04 24.63 25.44 28.20 30.60	3.6986 3.6115 3.4982 3.1617 2.9190	18.08 13.56 5.79 100.00 4.20	Integrated Intensity 150.875 5.790	m m m n	94.215
24.04 24.63 25.44 28.20 30.60 31.48	3.6986 3.6115 3.4982 3.1617 2.9190 2.8398	18.08 13.56 5.79 100.00 4.20 78.48	Integrated Intensity 150.875 5.790	m m m t	94.215

Cont	:d.	_	_	_	CE	3T	7

20 dogrees	d Å	I I _O	Integrated Intensity	Remarks	% phases
40.77	2.2112	24.77	en er	m	وي د ع
41.39	2.1796	14.47	وتري لاسيا	m	State 1 2
44.78	2.0222	12,66	gat.	m	هدره هدره
45.55	1.9897	_18:08	en 400	m	65.50 inv 1
49.30	1.8468	40.33	من ده	m	ಮೂ ಒತ
50.32	1.8117	48.82	కపడా	in	eg e-
50.50	1.8057	47.92	g) (t	Racion (CLL)
54.05	1.6951	23.51	No. Co.	m	ديون
55.47	1.6575	37.98	C.v.d stCts	m	هسه وليج
55.94	1.6423	25.32	819 tm	m	جنبة حثنه
57.22	1.6060	13.56	r.acii	m	بته بت
58.13	1.5855	16.28	دعده	m	ectis tomi
59.93	1.5421	34.72	الإيكا فسأله	m	ت ۵۰ فینت
61.35	1.5098	15.19	نيوب	m	د عدد ع
61.93	1.4970	16.28	ब्रह्म स्थाने	m	ويست فيسته
62,90	1.4763	20.43	Con con	m,t	6d 6 i
64.34	1.4467	7.41	क्षांत क्षण्य	m	*** #**
65.68	1.4204	19.89	engle leg ch	m	<u></u>
69.12	1.3578	6.69	ero esta	m	ه ۱۰ اهمیکا
21.21	1.3230	15.37	ಳು ಆತ	m	وساد سا
72.38	1.3045	6.87	_{கிறி} வீட 7	m	المسية المسية
75.25	1.2617	12.66	Carn	m	خيت وشرع
					The United States of the Control of

Table 4.4.h. -- Composition T7

20 degrees	d Å	I I _o	Integrated Intensity	Remarks	% phases
24.27	3.6641	12.41	ديثه	m	gaza t. "I
24.53	3.6113	10.86	600 600	m	سے بری
25.48	3.4928	3.62	ente ente	m	فنه دنو
28.30	3.1508	100.00	136.200	m	99.479
30.54	2.9283	0.90	1.143	m	0.521
31.60	2.8289	56.55	esc.	m	era e o
34.22	2.6166	22.40	523 (m.)	m	دے ہے
34.43	2,6098	24.48	£3 (-25	m	డుకి గునె
35.45	2.5300	18.10	هده کستا	m	cus cus
38.74	2.3224	9.48	65m.3 Ni .4	m	فيئ فسة
40.95	2.2071	18.97	627 tm3	m	بد) جي
41.60	2.1691	11.55	esse Cara	m	ھے بین
44.95	2.0148	9.83	هسه بسته	m	فسك هجيم
45.91	1.9955	12.41	co c	m	(mil 4+28
49.43	1.8422	13.69	యా కను	m	ಬ್ಯಾ ಭ೨
50.12	1.8185	36.38	ويبوع	m	euiza etaza
50.63	1.8013	38.45	au c	m	نستا حبيه
51.40	1.7762	15.17	ento Cus	m	e2 ##*
54.13	1.6929	15.52	තුරක වෙන	m	وتتاحب
55.54	1.6532	26.38	etten 4 va	m	80D) 90D

Contd....T7

Contd...T7

20 degrees	d A	I I ₀	Integrated Intensity	Remarks % p	hases
56.07	1.6388	20.00	دست دست	m ea	
57.34	1.6055	14.66	دسة دست	m	a s.a
58.38	1.5794	15.86	రా త రమ	m =	23 hand
60.18	1.5364	32.76	cm; sains	nı -	7 4. 3
61.41	1.5085	10.86	الله دسا	m	w =_3
61.90	1.4977	15.00	<u> ಅ</u> ವಾ ಒಡಕಿ	m	es em
62.98	1.4746	14.14	14.8 C 3	m -	a (-)
64.38	1.4459	6.55	ery cm	m	يت ۾
65.79	1.4182	18.45	دسا شا	m -	3 428
69.18	1.3568	4.83	ذنه ۱ درعه	m	= 43
71.20	1.3232	12.24	కుని గుమ	m	ø es
72.85	1.2972	5.17	600 feet	m -	اليكه جد
75.51	1.2580	12.24	دين فيته	m «	ے دین

Table 4 .4.i Composition M8					
20 degrees	d A	<u> </u>	Integrated Intensity	Remarks	% phases
24.05	3.6574	17.17	Alley 6-79	m	فيت فينا
					ಕ್ಷ-ಚಿಕ್ಕ ಿ
28.20	3.1617	100.00	इस्स १६७	Ţ	
31.30	2.8553	51.50	حة ب	m	دەن
33.99	2.6353	22,22		m	ئىن ئىن
35.15	2.5509	21.06	දෙය රැය	m	حيث فنية
38.57	2.3312	8.33	cups daile	m	ह्यूक स्थाप
40.78	2.2107	2.50	ಜನ ೯-೨	m	475 673
44.82	2.0204	15.15	සා දැය	m	and store
45.33	1.9988	18.20	هيئة جنة	m	ens and
49.11	1.8537	47.98	دينه هينا	m	೭.೨ ದನ
50.21	1.8155	67.42		m	613 62.4
53.93	1.6985	17.70	v ఎ కమ	m	فينة فين
54.33	1.6871	47.22	ಬ್	m	وساجب
57.10	1.6117	11.87	6.20 Ga2	m	دے ورے
58.00	1.5888	15.90	edis ciss	m	3 من ا الليبية 1 من ا
59.92	1.5424	37.52	sur ed	m	ند که هلیری
61.34	1.5100	18.94	pica c.a	m	සො ය
61.80	1.4999	23.20	GO L.X	m	هشه هشه
62.60	1.4827	18.94	ಕ್ಷವಾ ಕ್ಕುವಿ	m	دے دے
64.03	1.4529	3 9.60	అల	m	೯೫ ೯೨
65.50	1.3880	20.67	യ ം അ	m	(C.2)
69.01	1.3597	8.33	ಪ್ರಾಯಾ	m	දකු ය ා
71.10	1.3248	17.68	CENTRAL	THR AR	,
			The same appropriate in the same of the sa	A 6303	

Table 4.5.b. - Composition C13 Spalled

20 degrees	d A	I I _O	Integrated Intensity	Remarks % phase
24.4	3.6949	21.67	ONDERFORMATION AND A MALE AND A M	m
28.45	3.1345	100.00	65 . 0 5	m 87.483
30,42	2.93 5 8	16.67	14.92	c.t 12.517
31.70	2.8202	100.00		m
34.40	2.6048	32.92		m,t
35.43	2.5313	22,92		С
41.07	2.1958	22.50		c,t
45.03	2.0115	20.83		m
48.56	1.8732	42.92		m
50.40	1.8090	62.50		c,m,t
53.19	1.7205	28.33		t
55.60	1.6515	37.92		m
57.53	1.6006	28.33		m
58,30	1.5813	18.57		t,m
60.14	1.5373	41.67		c,t,m
62.68	1.4810	31.20		t,m
65.90	1.4161	29.17		m
71.52	1.3181	16.25		t,m

Table 4.5.c - Composition C10 Spalled

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% Phases
24.22	3.6716	16.80		m	
25.60	3.4767	4.56		m	
26.60	3.3982	1.82		m	
28.38	3.1421	100.00	108.575	m	76.64
30.33	2.9402	22.00	53.050	t	23.36
31.57	2.8315	93.41		m	
34.27	2.6137	41.82		m	
35.39	2.5,341	25.23		m	
38.72	2.3235	6.82		m	
41.10	2.1943	19.32		m	
41.38	2.1801	17.95		m	
44.92	2.0162	17.95		m	
45.50	1.9918	17.27		m	
49.60	1.8364	42.05		m,t	
50.32	1.8118	55.00		mt,	
54.20	1.6909	29.55		m	
55.60	1.6515	42.95		m	
57.25	1.6078	16.60		m	
58.48	1.5767	15.45		m	
60.13	1.5371	32.05		m	

Contd....C10 spalled

COTTOM S S S S C TO DOGT TEG	Con	td.			.C10	Spalled
------------------------------	-----	-----	--	--	------	---------

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% Phases
61.40	1.5087	22.73		m	an englist sell "Maller, anglis vert anglis betti, albut et aut a rei Spanis at a
61.88	1.4982	25.00		m,t	
62.78	1.4809	29.55		m,t	
64.34	1.4467	1.36		m	•
65,65	1.4310	22.50		m	
68.87	1.3621	7.50		m,t	
71.20	1.3232	16.14		m	
75.08	1.2641	10.90		m,t	

Tab.	Le	4.	5	.d	• «=	Com	pos.	iti	on	C7	Spal	led

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% Phases
24.23	3.6741	12.22		m	
25.45	3.4969	4.42		m	
28.35	3.1354	100.00	194.03	m	91.741
30.37	2.9406	8.84	28.00	t	8.259
31.60	2.8289	84.83		m	
34.32	2.6107	20.18		m,t	
35.40	2.5334	9.28		m	
38.73	2.3230	3.53		m	
40.92	2.2040	10.31		m	
44.98	2.0136	8.25		m	

Contd...C7 spalled

Contd	Ca	Spalled
COLLUGA		Sparred

20 degrees	d Å	IO	Integrated Intensity	Remarks	% Phases
45.40	1.9959	7.36	Process Transiscoppie, editore, Carloscoppie, apriliante, apri	m	
49.42	1.8426	18.70		m	
50.30	1.8124	23.86		m,t	
50.87	1.7934	22.09		m	
51.29	1.7797	8.84		m	
52.20	1.7508	2.65		m	
54.14	1.6926	10.01		m	
55.55	1.6526	14.58		m	
57.26	1,6075	7.07		m	
58.39	1.5791	7.36		m,t	
60.12	1.5379	14.43		m,t	
61.46	1.5074	6.19		m	
62.07	1.4940	6.33		m	
62.94	1.4755	8.98		m	
64.50	1.4434	3.53		m	
65.83	1.4369	9.28		m	
71.40	1.3200	4.70		m	
76.23	1.2479	3.40		m	reportioneres, angular a distributantan l'adicadendens

Table 4.5.e. - Composition Stab Z Spalled

20 degrees	d Å	<u> </u>	Integrated Intensity	Remarks	% Phases
24.17	3.6791	4.44		m	
28.31	3.1497	67.80	135.85	m	
30.23	3.9544	79.84	165.17	c,t	
31.50	2.8376	28.40		m	
33.26	2.6914	5.24		m	
34.20	2.6175	29.10		m	
35.22	2.5464	26.98		m,c	
36.86	2.4383	2.86		m	
38.82	2.3177	2.86		m	
40.83	2.2082	9.52		m	
45.57	1.9889	12.60		m	
49.32	1.8461	22.20		. m	
50.60	1.8024	100.00		t,c	
53.96	1.6987	7.30		m	
55.57	1.6524	18.40		m	
60.19	1.5362	61.90		t,c	
63.00	1.4742	17.46	· ·	m,c,t	
71.35	1.3208	6.98		m	
74.47	1.2730	6.35		m,t,c	

Table 4.5.f	Composition	C7T5	Spalled

20 degrees	d Å	<u>I</u>	Integrated Intensity	Remarks	% Phases
24.34	3.6527	17.68		m	
28.40	3.1399	100.00	88.93	I U	80.779
30.40	2.9377	24.39	33.92	t	19.221
30.95	2.8869	14.33		m	
31.62	2.8272	87.20		m	
34.37	2.6070	47.56		t,m	
35.46	2.5293	24.39		t,m	
38.80	2.3189	6.71		m	
41.00	2.1994	22.87		m	
41.36	2.1811	22.87		m	
45.20	2.0043	21.65		m	
45.62	1.9465	12.20		m	
49.50	1.8398	45.73		t,m	
50.34	1.8110	58.23		m	
50.63	1.8014	58.84		m,t	
54.28	1.6885	30.49		m,t	
55.63	1.6507	50.61		m	
57.27	1.6073	10.37		m	
58.46	1.5772	11.28		m	
60.15	1.5371	36.59		m,t	
61.18	1.5137	18,90		m	

Contd....

Contd....C7T5 Spalled

20 degrees	d Å	I	Integrated Intensity	Remarks	% Phases
62.07	1.4940	21.95	and the second s	m	ONLINE : 1 MILES MICORDIL COMES COME
63.03	1.4736	18.90		m	
64.53	1.4428	6.71		ih	
65.87	1.4361	21.65		m	
71.34	1.3194	19.82		m	
	Table ·	4.5.g	Composition C8	T7 Spalled	ng ang panggang at mangkang panggang ang panggang ang panggang panggang panggang panggang panggang panggang pa
20 degrees	å Å	I	Integrated Intensity	Remarks	% Phases
24.49	3.6317	18.67		m	
25.83	3.4463	5.90		m	
28.62	3.1163	95.82	121.10	m	98.97
30.55	2.9237	3.50	2.06	t	1.03
31.81	2.8107	100.00		m	
34.45	2.6011	39.31		m	
36.62	2.4518	16.71		m	
39.02	2.3063	7.37		m	
41.18	2.1902	25.31		m	
45.12	2.0119	16.22		m	
45.81	1.9791	22.10		m	
46.27	1.9604	4.18		m	

Contd.....

Contd....C8T7 Spalled

20 degrees	d A	I I _O	Integrated Intensity	Remarks	% Phases
49.67	1.8340	55.04	от температично при	m	
50,52	1.8051	56.51		m	
54.34	1.6868	33.91		m	
55.76	1.6472	41.77		m	
57.53	1.6006	20.39		m	
58.40	1.5789	17.69		m	
60.27	1.5343	28.99		m	
61.73	1.5014	16.95		m	
62.15	1.4923	19.66		m	
63.03	1.4736	26.54		m	
64.55	1.4424	11.79		m	
66.11	1.4121	29.48		m	
71.60	1.3167	19.90		m	
72.53	1.3022	12.53		m	

Table 4.5h-- Composition T7 Spalled

grees	d Å	I I _O	Integrated Intensity	Remarks	% Phases
,24	3,6686	11.67		m	
,58	3.6186	12.92		m	
43	3.1367	66,67		m	
.60	2.8289	100.00		m	
. 28	2.6130	18.54		m	
. 57	2.5217	27.50		m	
. 82	2.3177	7.50		m	
.21	2.1938	18.75		m	
•98	2.0136	15.83		m	
.78	1.9103	12.71		m	
. 50	1.8393	29.17		m	
.51	1.8054	47.08		m	
.32	1.7788	45.00		m	,
.73	1.6480	33.33		m	
•23	1.6083	18.33		m	
.12	1.5377	21.04		m	
• 59	1.5045	16.04		m	
.12	1.4930	11.25		m	
.11	1.4719	33.33		m	
.99	1.4145	24,38		m	

Contd....T7

Contd. T7 Sp					
20 degrees	d Å	I I _O	Integrated Intensity	Remarks	% Phascs
69.39	1.3532	4.38		m	
71.48	1.3187	7.71		m	
72.79	1.2982	6.25	,	m	
73.60	1.2859	3.13		m	
	Table	4.5.i-Conp	ositions M8 S	palled	ngy yadhaddililaindac.um _{a.} yandi.ggung T937 I saddilinud. B
20 degrees	đ Å	I	Integrated Intensity	Remarks	% Phases
24.13	3.6851	14.34		m	
24.51	3.6288	13.44		m	
25.55	3,4834	4.12		m	
28.27	3.1541	100.00		m	
31.55	2.8333	79.75		m	
32.34	2,7658	28.32	•	m	
35.44	2.5306	19.71		m	
36.87	2,4357	6.81		m	
38.69	2.3253	5.38		m	
40.85	2.2071	19.71		m	
44.98	2.0136	12.00		m	
45.62	1.9868	10.75		m	
49.40	1.8433	23.30		m	
50.40	1.8124	32.26	•	m	

Contd....M8

td... M8 Spalled

degrees	d A	I I _O	Integrated Intensity	Remarks	% Phases
•09	1.7235	15.77	M. A. Malay - announce and a finished at the lead of the shadow, and a supply of Discourage of America	m	AND THE PROPERTY OF THE PROPER
.48	1.6549	24.37		m	
.33	1.6057	8.24		m	
.28	1.5818	7.17		m	
.30	1.5336	16.70		m	
•97	1.4748	16.13		m	
.80	1.4180	15.23		m	

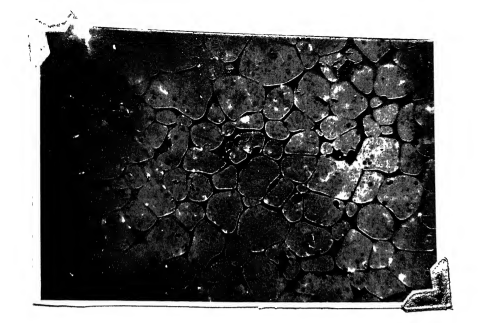
phase field. So some amount of tetragonal form was produced which during cooling got transformed to the monoclinic polymorph.

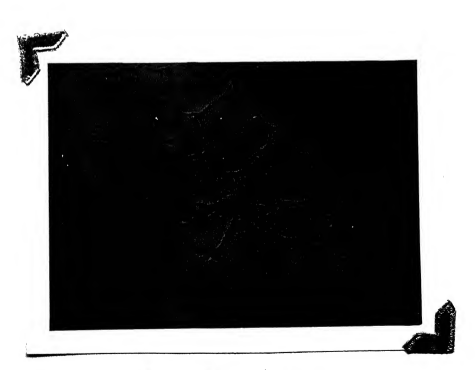
4.3 Microstructure by optical microscopy

Grain size, grain size distribution, porosity, etc. for sintered specimens have been studied by optical microscopy, figures 4.9(i)-4.9(ii)The average grain size for the batches C7, C10, C13, C15 & Stab Z are 14.3., 7.2 , 10, 13.9 ,, 18.6 microns respectively. It has been found that except for the batch C7, the grain sizes of the other calcia-zirconia batches have increased with increasing CaO content. This can be explained by the increased diffusion rate due to larger number of vacancies for higher CaO-containing batches. Though the grain size of the C7 batch is high, yet there are enough intergranular and intragranular pores to make the density low. With the addition of TiO, to CaO-ZrO, grain size has decreased as C7T5 batch has an average grain size of 11.4 micron and for C8T7 batch the grain size is 4.3. micron. The reason lies in the lower diffusion rate due to lower vacancy concentration.

4.4 Vickers Hardness

Vickers hardness numbers (VHN) are shown in table 4.6 and fig. 4.10. It has been found that for CaO-ZrO2 systems





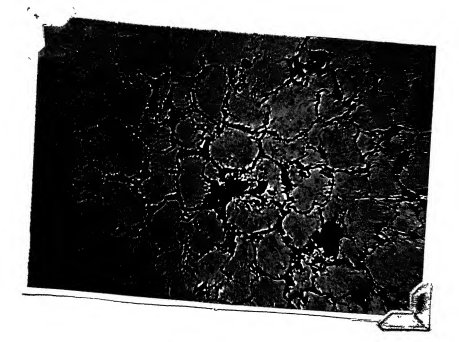
9.5. 00. (ii) St-63



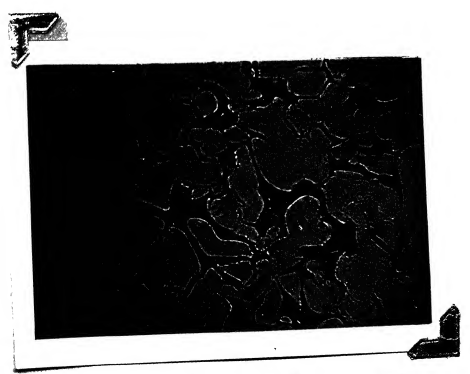
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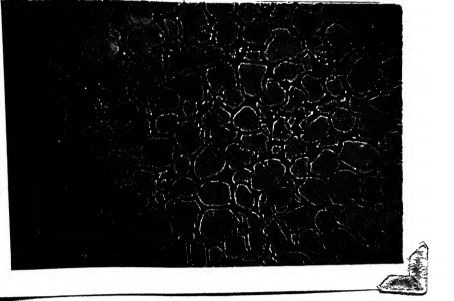
4 ig 21.7.(iv)



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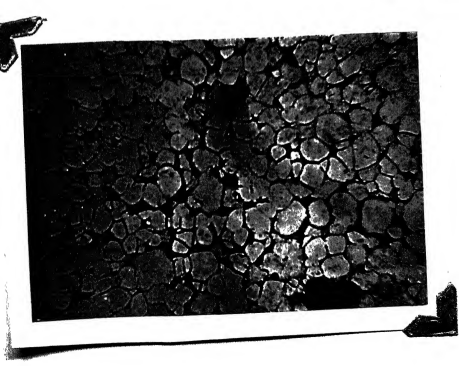


Rig. 48 (VI) M8



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TABLE 4.6

VICKERS HARDNESS & INDENTATION CRACK LENGTHS

Compositon L	oad in Kg 2	Length of diagonal x 10 ⁻³ mm	Crack length x 10 ⁻³ mm	VHN
1	antia particologica de la compania del la compania de la compania del la compania de la compania del la compania de la compani	3	Λ	
AND DESCRIPTION OF THE PROPERTY OF THE PROPERT	2 E		4	5
Stab Z	2.5	63	94	1168
	2.5	56	71	1478
	2.5	55	63	1533
	5.0	82	95	1379
	5.0	86	54	1253
C10	2.5	90	No crack	572
	2.5	69	ti	974
	2.5	83	и	673
	2.5	91	11	560
	5.0	126	25	584
C13	2:5	75	No erael:	824
	2.5	85	tt.	641
	2.5	73	и .	871
	2.5	70	ts	946
	2.5	70	u	946
	5.0	123	18	613

Contd...

Contd...4.6

1	2	3	4	5
C15	2.5	75	86	824
	2.5	75	93	824
	2.5	7 9	147	743
	2.5	69	167	976
	5.0	129	55	558
	5.0	133	7 9	613
C7T5	2.5	86	No crack	626
	2.5	88	tt	598
	2.5	82	tt	689
	2.5	88	13, Very thin crack	598
	5.0	125	No crack	595
C8T7	2.5	81	No crack	706
	2.5	86	u	626
	2.5	82	u	689
	2.5	85	и	641
	2.5	82	н	689
	5.0	126	16	593
	5.0	114	14	713

Contd.6

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Contd... 4.6

AND AND ADDRESS OF THE PARTY OF		Manager of the Commence of the		
1	2	3	4	5
C7	2.5	87	No crack	612
	2.5	91	13	560
	2.5	73	18	870
	2.5	90	23	572
	5.0	145	No crack	441
т7	2.5	100	18	463
	2.5	93	12	536
	2.5	92	13	547
	2.5	93	15	536
	5.0	140	83	473
	5.0	288	30	112
	5.0	181	13	283

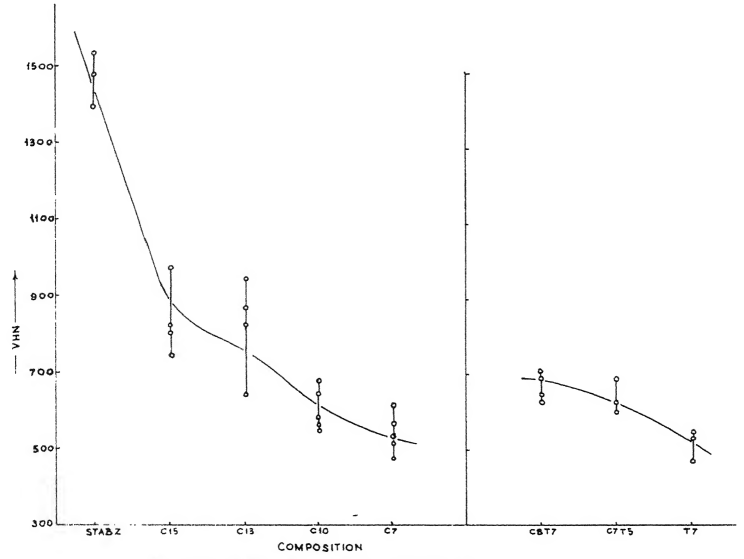


FIG 410 VICKERS HARDNESS NO VS. COMPOSITION

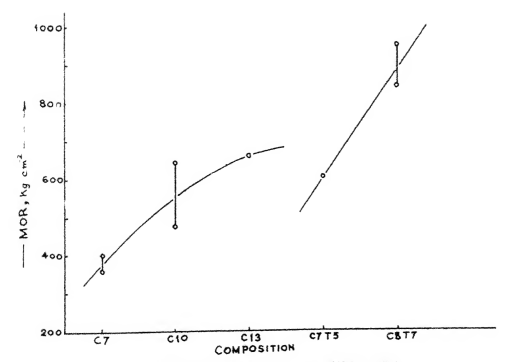


FIG. 4-12 MODULAS OF RUPTURE VS. COMPOSITION.

VIIN has increased with increase of CaO content which is natural as the higher CaO containing batches are of higher density. For the CaO-TiO₂-ZrO₂ batches the trend is similar i.e., with increase of density the VHN has increased.

4.5 Fracture toughness

The fracture toughness values of different batches as determined by indentation technique are shown in table 4.7. The values were obtained by first estimating the quantity

$$\frac{\Phi}{H\sqrt{a}} \left[\frac{H}{\Phi E} \right]^{0.4}$$

by using Young's modulus E ≈ 155.7 GN/m² for C7, C10, C8T7, C7T5 and T7, 80 GN/m² for C13 and 67.53 GN/m² for C15 and Stab Z (35). The constraint factor was taken as 3 (34). Hardness, H was obtained from equation 3.12 (33) using the experimental values of impression radius a. The dimensionless parameter $K_{C} \longrightarrow H_{C} \longrightarrow H_{$

The C15 composition has the lowest fracture toughness, Stab Z has similar value. The fracture toughness values of C7, C10, C13 are much higher than those of C15 and Stab Z

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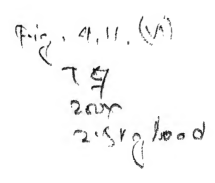
215 kg land

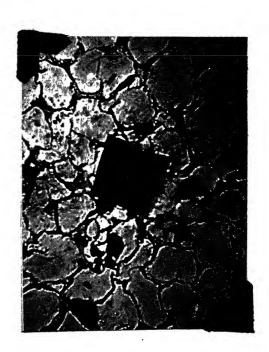
1.5. 4.11. (111)
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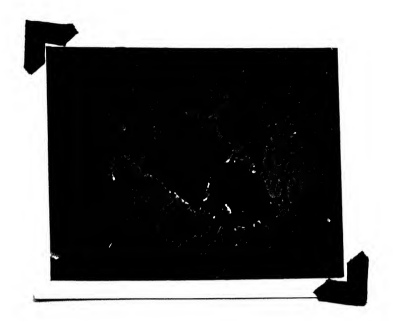




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TABLE 4.7
FRACTURE TOUGHNESS

Composition	Impression radius a x 10 ⁻⁶ m	Average crack length C x 10 ⁻⁶	<u>C</u> a dimen- sion- less	H -2 GNm	E -2	$\phi(\underline{H})^{0.4}$ $\zeta^{\prime,E}_{\text{GN}} 1_{\text{m}} 3/2$	kc(H) CEE Hadimension less	0.4 KC 3/2
1	2	3	4	5	6	7	8	9
Stab Z	86	54.5	0.6337	14,484	67.53	9.7666×10	-3 _{0.054} 0	5.527
	56	71.0	1.267			9.6349x10	-3 _{0.045}	2 4.691
	55	63.0	1.146			9.7220x10	⁻³ 0.047:	1_4.840
والمراور والاراد المراد	وفيينا حجو عوال كالمراجو المراجو الموري	23 6 4 4 4 4 4 4 4 4 4 4 5 4 4 7 79 6	ے دی جہ دے جہ دے دے دے	دے دیا ہے تک لیک کیا دے	د، شاهه دسانب کتار	على والما والما والما والما والما والما والما والما		سام در ۱ است السنا السنا
C15	75	93.0	1.240			1.163x10		
	75	86.0	1.147	8.295	67.53	1.1630x10	-2 _{0.047}	1 4.045
	79	147.0	1.861			1.1333x10	⁻² 0.037	1 3.269
	128	78.0	0.609			8.9033x10	⁻³ 0.05 5	3 6.205
C13	123	18.0	0.1463	8.075	:80.00	6.6087x10	-3 _{0.059}	0 8.626
C10	126	25.0	0.1984	6.142	155.7	6.529x10	3 0.059	3 7.691
C7	91	13.0	0.143	23		9.840x10	3 0.059	8 6.072
	145	5.0	0.007	5.367	155.7	7.80×10^{-3}	0.060	7 8.076
	90	23.0	0.256			9.87x10 ⁻³	0.063	5.997

Contd....

1	2	3	4	5	6	7	8	9
C8T7	126 114	16 14	0.127 6 0.123	. 755	366 7	7.2677x10 ⁻³		8.290 7.893
C7T5	86 82	1	0.023 6 0.012	.142	155 7	9.3137x10 ⁻³ 9.538x10 ⁻³		6.678 6.521
T7	140 181 93 100 92	53 13 12 18 13	0.3786 0.0718 5 0.129 0.18 0.1413	.252	155.7	8.0186×10 ⁻³ 7.0522×10 ⁻³ 9.8380×10 ⁻³ 9.488×10 ⁻³ 9.892×10 ⁻³	0.0617 0.0602 0.0598	7.172 8.749 6.119 6.303 6.040
M8	383 218 209	110 146 149	0.287 0.6697 0.713	5.4	155.7	4.7678x10 6.3196x10 6.4543x10	0.0468	7.406

as shown in the table.

With addition of TiO₂, in all the cases, the fracture toughness values have increased. T7 has the highest fracture toughness. C7T5 and C8T7 are also good in respect of fracture toughness.

In C7, C10, and C13 batches the major phase is monoclinic and the rest is tetragonal for C7, C10 and cubic for C13. But C15 and Stab Z are nearly single phase cubic zirconia. It isawell known fact that the fracture toughness value of cubic ZrO_2 is low which explains the lower fracture toughness of C15 and Stab Z batches.

form

C7T5, T7 and M8 batches/fully monoclinic solid solutions on sintering. C8T7 in addition to being a monoclinic SS, contains a small amount of tetrogonal ZrO2.

The tougher materials at the same time are of low hardness value. This can be explained from the equation number 3.10. Lower hardness means higher indentation impression radius (a) and consequently higher $K_{\mathbb{C}}$ or in other words higher toughness.

The low fracture toughness value in C15 and Stab Z is evident from the figure (4.11a-i) showing the indentations surrounded by many big cracks even with 2.5 Kg load. Indentation in T7 developed small cracks with 2.5 kg load. Rest of the

batches did not show any crack at 2.5 kg load. The later batches excepting C7 however developed small cracks at 5 kg load.

Finally, it will be worthwhile to mention here that our fracture toughness values are comparable or even better than the values obtained by other workers. This has been shown in table numbers (4.7 & 4.8).

TABLE 4.8
FRACTURE TOUGHNESS VALUES OF DIFFERENT MATERIALS

Composition	K _C MN m ^{-3/2}	Reference
Mgpsz	26	Porter and Hener (12-22)
Y203psz	6-9	Gupta, Lange etal (13)
BC	6	Evans & Charles (32)
Saphire	2.1	44 11
Spinel	1.31	g tr
Si ₃ N ₄	4.9	n n
SiC	4.	н н

Table 4.9 Modulus of rupture

mposition	Thickness of the sample d,mm	Width of the sample b,mm	Load, Kg	MOR 2 Kg/cm.	MOR, x10 ³ psi
T5	8.2	5.62	30	608.13	8.65
г7	7.9	5.46	42.9	956.33	13.6
				843.27	11.99
				847.65	12.10
3	7.92	5.60	31.3	661.60	9.41
0	8.54	6.60	45.50	642.13	9.13
				473.30	6.73
	8.95	6.00	12.60	400.24	5.69
				356.92	5.07
	9.83	5.98	31.7	473.44	6.73
				736.06	10.47

4.6 Modulus of Rupture (MOR)

MOR values are recorded in table 4.9 and the variation of MOR with composition is shown in the figure 4.12. It is evident from the table that with addition of TiO₂ the MOR value has increased. As for example, the MOR value of C7 batch is nearly 375.Kg/cm² and for C7T5, it is 608 Kg/cm². One of the possible reasons is the higher density of the TiO₂ added batches. The other reason may be in higher toughness value in TiO₂ containing batches.

4.7 Thermal Shock Resistance

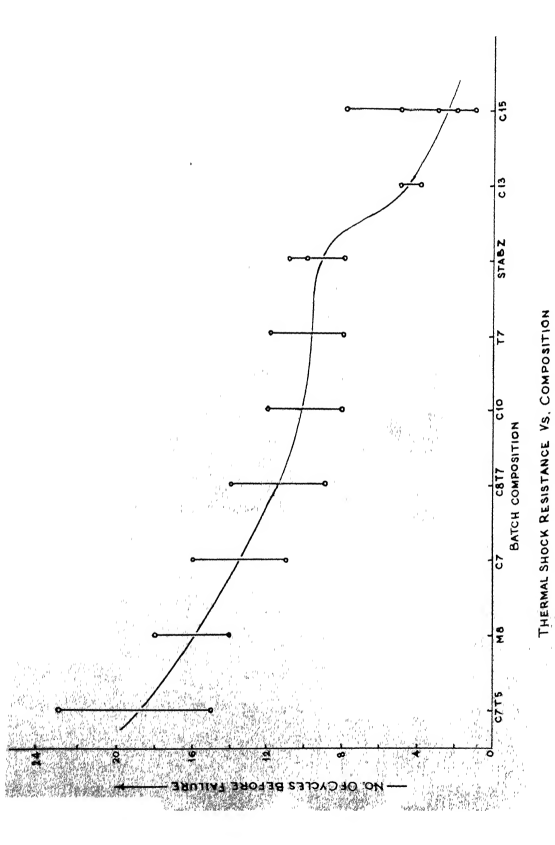
Thermal shock resistance is measured by the number of thermal cycles excluding the last cycle, the last cycle being defined by the chipping, or crumbling of the samples, or pulling apart by tongs. Figure 4.13 shows the number of cycles before failure as a function of composition. This figure gives an obvious trend of decreasing spalling resistance from C7T5 to C15 in the following order C7T5 M8 C7 C8T7 C10 T7 Stab Z C13 C15.

The Cl3, Cl5 and Stab Z batches contain higher percentage of cubic phase and have much lesser thermal shock resistance than the others. The C7T5 batch has the highest thermal shock resistance.

TABLE 4.10
THERMAL SHOCK RESISTANCE

Batch	No. of Cycles before failure	Batch	No. of Cycles before failure
C7	16	С7Т5	23
C7	11	C7T5	15
C10	12	C8T7	14
C10	8	C8T7	9
C13	4	M8	18
C13	5	M8	14
C15	5	M8	14
C15	5	т7	12
C15	8	т7	8
C15	3	т7	8
C15	1	Stab z bushing	10
C15	2	Stab z	11
C15	3	bushing	11
		Stab z bushing	8

C7T5 > M8 > C7 > C8T7 > C10 > T7 > Stab z > C13 > C15



Thermal shock resistance (spalling resistance) is a property which is dependent on many factors namely thermal conductivity, thermal expansion coefficient, temperature gradient, grain size, grain size distribution, pore size, pore size distribution, presence of glassy phase, type of phase of $2rO_2$, etc. So more detailed work is required to find out the possible reasons of the above trend in thermal shock resistance.

4.8 Corrosion resistance

The corrosion resistance of the zirconia crucibles in contact with E glass has been found to be good. The crucibles containing the E glass in the corrosion resistance test did not get affected, except only a little thinning at the airglass-2rO₂ interface for some batches. TiO₂ containing batches also showed similar corrosion resistance properties. The interface corrosion is the least in C7 crucibles. Next is M8 crucible. But corrosion in the bulk is not observable in any of the batches. In contrast, the Al₂O₃ crucible has got heavily corroded; Corrosion at the air-glass Al₂O₃ interface is more pronounced than in the interior. In case of Al₂O₃, a layer of intermediate compositions between Al₂O₃ and glass has formed by dissolution of Al₂O₃ which rendered it visible. The photographs of the crucibles after corrosion resistance test are shown in fig. 4.14.

4.9 Discussions on experimental procedure

4.9.1 Grinding

Crinding was done in alumina lined ball mill and agate ball mill. In order to avoid extraneous contamination from the jar and balls; grinding could be done in steel jars using steel balls; and then the incoming iron has to be removed by magnetic separation. Longer grinding of C8T7 and C7T5 batches did not produce remarkable effect on the particle size distribution as compared to CaO-ZrO₂ batches because of the fact that these two batches got calcined to a greater strength than the others. So, in general, C8T7, C7T5 batches seem to need longer grinding.

4.9.2 Pressing

A moderately high pressure was used. Very high pressure causes steep pressure gradient contours.

4.9.3 Sintering

We used 1 hour soaking period at 1950°C, but longer period can be used for higher sintered density. More work is required to optimise the soaking period & temperature.

The zirconia batches should be tried for further heat treatments of solution annealing at 1800-1900°C for dissolving the monoclinic phase, quenching to 1000°C followed by ageing

etween 1250 to 1350°C and finally slow cooling. This may

4.9.4 X-ray diffraction study

It was done with as sintered bars. It was not ground to powder as grinding and mechanical stresses trigger tetragonal-monoclinic phase transformation and leads to wrong idea about the tetragonal phase (22, 24) by X-ray diffraction.

4.9.5 Thermal shock resistance

According to ISI Specification (36), the temperature used for thermal cycling to measure the thermal shock resistance for SiO₂ bricks is 450°C, and that for fire brick, siliceous and basic bricks, is 1000°C. Since ZrO₂ has much higher melting temperature (2690°C), so we have used 1300°C for thermal cycling.

4.9.6 Corrosion resistance.

More work is required to establish the corrosion resistance property of our samples. We have studied the corrosion resistance of our samples in contact with E glass (which is more corrosive than A, C, S and HS glasses). But corrosive effects of other glasses and slags are to be tested. At the same time, long duration corrosion resistance tests are also necessary to characterize these materials properly.

4.9.7 Mechanical Properties

Though we have measured MOR, VHN and Kc, but young's modulus in tension should also be measured for better characterization of our materials:

CHAPTER 5

CONCLUSION

- 1. The average particle size of the powder (~'0.2 micron) which we used for final sintering is optimum size, because higher particle size means settling of the slip used for slip casting and less densification and finer particles will give rise to drying cracks after casting.
- 2. With addition of ${\rm TiO}_2$ in ${\rm CaO-ZrO}_2$ systems, the bulk density has decreased at the cost of improvement of other properties.
- 3. In all the TiO_2 containing batches, the major phase is monoclinic, no cubic phase (ZrO_2) is formed. In C8T7, small amount of tetragonal phase is formed, whereas cubic phase is present in C13 and C15 batches, but no tetragonal phase has been found.
- 4. Grain size can be controlled to some extent by adding TiO2. Finest grain size has been obtained in CST7 composition.
- 5. With addition of CaO, the hardness (VHN) has increased in all the cases, the same is true for TiO₂ addition but the imcrease in hardness is not so appreciable.
- 6. For the same amount of CaO, MOR values have been increased on addition of ${\rm TiO}_2$. Maximum MOR has been obtained for C8T7 composition.

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